Introduction

Influenza A viruses (IAV) infect cells in the alveolus and induce primary viral pneumonia, which can progress to acute respiratory distress syndrome (ARDS) with high mortality (1). IAV-induced lung injury is characterized by exaggerated inflammatory responses and loss of barrier function, resulting in edema formation and severe hypoxemia (1–4). Resolution of inflammation together with repair of the damaged epithelium and edema fluid clearance are crucial events in the recovery phase of ARDS (5). It has been demonstrated that impaired edema resolution in ARDS patients is strongly correlated with increased mortality (6, 7). Therefore, understanding the molecular mechanisms by which IAV infection decreases lung edema clearance might provide therapeutic targets to improve clinical outcome in IAV-induced ARDS.

Fluid reabsorption is driven by an osmotic gradient generated by the coordinated regulation of the apically expressed epithelial sodium channel (ENaC) and the basolaterally expressed Na,K-ATPase. Sodium ions enter the epithelial cells via ENaC and are transported out of cells by the Na,K-ATPase, which promotes the intracellular transfer of 2 potassium ions with energy provided by the hydrolysis of ATP. It is therefore the major driver of vectorial sodium transport and has been shown to be an important determinant and limiting factor of net fluid clearance (10–15).

During IAV infection of the distal respiratory tract, the functional and structural impairment of the tight alveolar epithelial cell (AEC) layer, and thus the alveolo-capillary barrier results in fluid leakage from the vascular compartment into the alveolar space and persistent edema (4, 16, 17). However, the underlying mechanisms are not well understood. Recent findings suggest that an overly exuberant inflammatory response may contribute to alveolar barrier disruption via a mechanism that requires both tissue-resident and BM-derived macrophages (BMM) (4, 16, 18, 19). We have recently demonstrated that the cytokine TNF-related apoptosis–inducing ligand (TRAIL), when released in excessive amounts from aveolar macrophages (AMs) upon autocrine IFNβ stimulation, significantly contributes to IAV-induced immunopathology (4, 16).

Here, we identify a paracrine communication network between different cellular subsets within the inflamed alveolus, which we found is critical to affect alveolar fluid clearance (AFC) in IAV-induced lung injury. In vitro and in vivo IAV infection led to alveolar macrophages (AMs) upon autocrine IFNβ stimulation, significantly contributes to IAV-induced immunopathology (4, 16).

Macrophage–epithelial paracrine crosstalk inhibits lung edema clearance during influenza infection

Christin Peteranderl,1,2 Luisa Morales-Nebreda,3 Balachandar Selvakumar,1,2 Emilia Lecuona,1 István Vadász,1,2 Rory E. Morty,1,2,4 Carole Schmoldt,1,2 Julia Bespalowa,1,2 Thorsten Wolff,1 Stephan Pleschka,1 Konstantin Mayer,1,2 Stefan Gattenloehner,7 Ludger Fink,2,8 Juergen Lohmeyer,1,2 Werner Seeger,1,2,4 Jacob I. Sznajder,3 Gökhan M. Mutlu,9 G.R. Scott Budinger,3 and Susanne Herold1,2

1Department of Internal Medicine II, University of Giessen and Marburg Lung Center (UGMLC) Giessen, Germany. 2German Center for Lung Research (DZL). 3Division of Pulmonary and Critical Care Medicine, Northwestern University Feinberg School of Medicine, Chicago, Illinois, USA. 4Department of Lung Development and Remodeling, Max Planck Institute for Heart and Lung Research, Bad Nauheim, Germany. 5Division of Influenza Viruses and Other Respiratory Viruses, Robert Koch Institut, Berlin, Germany. 6Institute of Medical Virology and 7Department of Pathology, Justus Liebig University Giessen, Giessen, Germany. 8Institute of Pathology and Cytology, Wetzlar, Germany. 9Section of Pulmonary and Critical Care Medicine, The University of Chicago, Chicago, Illinois, USA.

Influenza A viruses (IAV) can cause lung injury and acute respiratory distress syndrome (ARDS), which is characterized by accumulation of excessive fluid (edema) in the alveolar airspaces and leads to hypoxemia and death if not corrected. Clearance of excess edema fluid is driven mostly by the alveolar epithelial Na,K-ATPase and is crucial for survival of patients with ARDS. We therefore investigated whether IAV infection alters Na,K-ATPase expression and function in alveolar epithelial cells (AECs) and the ability of the lung to clear edema. IAV infection reduced Na,K-ATPase in the plasma membrane of human and murine AECs and in distal lung epithelium of infected mice. Moreover, induced Na,K-ATPase improved alveolar fluid clearance (AFC) in IAV-infected mice. We identified a paracrine cell communication network between infected and noninfected AECs and alveolar macropages that leads to decreased alveolar epithelial Na,K-ATPase function and plasma membrane abundance and inhibition of AFC. We determined that the IAV-induced reduction of Na,K-ATPase is mediated by a host signaling pathway that involves epithelial type I IFN and an IFN-dependent elevation of macrophage TNF-related apoptosis–inducing ligand (TRAIL). Our data reveal that interruption of this cellular crosstalk improves edema resolution, which is of biologic and clinical importance to patients with IAV-induced lung injury.

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to reduced plasma membrane expression of alveolar epithelial Na,K-ATPase and thus impaired lung edema clearance in vivo. We discovered that IAV-infected epithelial cells and macrophages, as well as noninfected neighboring epithelial cells, establish a specific communication network via epithelial type I IFN and especially the IFN-dependent, macrophage-expressed TRAIL, which determines Na,K-ATPase plasma membrane protein abundance and, thus, edema clearance during IAV infection. The identified pathways underlying this macrophage-epithelial crosstalk are amenable to therapeutic targeting to improve alveolar edema clearance in vivo and putatively improve outcome in patients with severe IAV-induced lung injury.

Results

IAV infection leads to decreased Na,K-ATPase protein abundance and impaired AFC. Murine intratracheal infection with H1N1 IAV A/PR/8/34 resulted in alveolar edema at day 7 postinfection (d7 pi) not present in PBS-treated control groups (Figure 1A, upper panel). The excessive fluid in the alveolar airspace upon IAV infection was associated with severe hypoxemia at d7 pi (Figure 1B) and substantial impairment of AFC in vivo detected at d2 and more pronounced at d7 pi (Figure 1C). We therefore assessed Na,K-ATPase α1 subunit (NKAα1) protein abundance, after in vitro H1N1 IAV infection of primary AEC. Gene expression levels of NKAα1 did not change after IAV infection (data not shown); however, the total cell protein abundance was significantly decreased at 16 hours pi (16h pi) and 24h pi in murine AEC (mAEC) as well as in in primary human AEC (hAEC) at 16h pi (Figure 1, E and F). In addition, IAV infection was associated with reduction of the immunofluorescent signal for NKAα1 in vivo, which was particularly found within highly inflamed alveolar regions but was found less within less inflamed alveolar regions; noninfected mice showed high NKAα1 expression (Figure 1A, lower panels).

Alveolar epithelial plasma membrane NKAα1 is decreased after IAV infection in vitro. Na,K-ATPase is mostly stored in intracellular compartments but contributes to active ion transport and AFC when recruited to the basolateral cell membrane (20, 21). Therefore, we assessed NKAα1 plasma membrane abundance by flow cytometry (gating strategy in Figure 1D), as well as by plasma membrane biotinylation and subsequent pulldown in in vitro–infected AEC. Both assays revealed reduced NKAα1 protein abundance at the plasma membrane in mAEC and hAEC following IAV infection (Figure 1, G–I).

Reduced alveolar epithelial plasma membrane expression of NKAα1 causes impaired AFC after IAV infection in vivo. We next investigated whether Na,K-ATPase levels were affected by IAV infection in vivo and assessed NKAα1 plasma membrane protein abundance on distal lung epithelial cells, largely comprising type II alveolar epithelial cells (Supplemental Figure 1; supplemental material available online with this article; doi:10.1172/JCI83931DS1), by flow cytometry. IAV infection resulted in a reduction of plasma membrane NKAα1 on alveolar epithelial cells in vivo already on d2 pi and most pronounced on d7 pi (Figure 2A), which correlated with the reduced fluid clearance depicted in vivo (Figure 1C). To assess if decreased levels of NKAα1 were a direct cause of reduced AFC capacities after IAV infection, we either induced overexpression of NKAα1 and NKAβ1 (22, 23) in parallel by adenoviral delivery prior to IAV infection or increased Na,K-ATPase plasma membrane abundance by treatment of AEC by the β-adrenergic agonist salbutamol (15, 24). Adenoviral overexpression of NKAα1 and NKAβ1 indeed restored total NKAα1 in IAV-infected AEC to baseline levels in vitro (Figure 2C) and also plasma membrane expressed NKAα1 in vivo (Figure 2D). Similarly, salbutamol treatment increased NKAα1 plasma membrane abundance both in vitro and in vivo (Figure 2, B and D). Importantly, increased NKAα1 expression on alveolar epithelial cells directly contributed to improved AFC after IAV infection in vivo (Figure 2E). In line, direct blockade of Na,K-ATPase activity by its specific inhibitor ouabain resulted in decreased AFC (Supplemental Figure 2).

A soluble mediator released from infected macrophages and AEC leads to decreased plasma membrane NKAα1 protein abundance. Given our previous findings that alveolar and BMM contribute to AEC damage (4, 16) and that NKAα1 was particularly reduced within strongly inflamed regions of the lung, we explored whether presence of AMs or BMM would further inhibit NKAα1 after IAV infection. Indeed, epithelial cell NKAα1 protein abundance of noninfected AEC was significantly reduced in the presence of infected AM alone and further reduced after infection of both AEC and either AM (Figure 3A) or BMM (Figure 3B). Notably, plasma membrane–expressed NKAα1 was significantly decreased on noninfected AEC in presence of IAV-infected BMM (Figure 3, C and D). As AEC and AM/BMM were cultured without direct physical contact to each other, these results suggest that paracrine crosstalk via a soluble macrophage-produced mediator leads to NKAα1 downregulation and that additional infection of AFC further reduced total and plasma membrane–expressed NKAα1 (Figure 3, C and D). We therefore investigated whether the reduction of NKAα1 at the plasma membrane after IAV infection in the AEC monoculture was similarly mediated by paracrine signaling rather than by IAV infection itself. We correlated viral protein expression (hemagglutinin, HA) to NKAα1 plasma membrane abundance (Figure 3E), and indeed, plasma membrane NKAα1 was predominantly reduced in the adjacent noninfected IAV-HA fraction of mAEC (Figure 3F) and hAEC (Figure 3G), as well as of alveolar cells after IAV infection in vivo (Figure 3H). Furthermore, treatment of mAEC with conditioned media of infected but not of PBS-treated mAEC was sufficient to decrease NKAα1 plasma membrane protein abundance (Figure 3I).

Paracrine signaling of epithelial type I IFN- and IFN-dependen macrophage TRAIL leads to reduced plasma membrane NKAα1 protein abundance in AEC. To identify soluble factors within the AEC-macrophage crosstalk network that mediated the observed effect on NKAα1 plasma membrane expression in AECs, we analyzed coculture supernatants for proinflammatory cytokines by ELISA and cytometric bead array (data not shown). We found that IFNα — and to a minor extent IFNβ (Figure 4A and Supplemental Figure 3) — were released from infected AEC, whereas BMM did not increase IFNα release upon IAV infection (Figure 4A).

We recently demonstrated that type I IFNs are potent inducers of TNF family cytokines in AM, particularly of TRAIL (4, 16). Likewise, IAV infection of BMM with IAV led to a significant release of TRAIL (Figure 4B), which was type I IFN–dependent as demonstrated by use of BMM derived from type I IFN receptor–deficient
dance was rescued to baseline levels at 24h pi, whereas Trail–/– AEC still displayed a strong decrease in NKAα1 protein abundance (Figure 4E). Moreover, transfer of conditioned media from either PBS-treated or PR8-infected AEC to Ifnar–/– AEC did not reduce NKAα1 plasma membrane abundance (Figure 4F), suggesting that in absence of macrophages, NKAα1 surface downregulation was solely dependent on signaling through type I IFN. We next performed coculture infections using AEC and BMM from WT, Ifnar–/–, Trail–/–, or Dr5–/– (TRAIL receptor) mice and assessed NKAα1 plasma membrane abundance after IAV infection. These (Ifnar–/–) mice (Figure 4C). AEC did not release TRAIL in response to IAV infection (Figure 4B). Treatment of uninfected AEC with either recombinant IFNα (rIFNα) or rTRAIL decreased NKAα1 protein abundance at the plasma membrane, and combined treatment with both IFNα and TRAIL reduced NKAα1 to similar levels as coinfection of AEC and BMM (Figure 4D). rIFNβ at levels found when released by infected AEC did not affect NKAα1 levels (Supplemental Figure 3). When using Ifnar–/– AEC to block paracrine IFNα signaling between A/PR/8/34 (H1N1)–infected (PR8-infected) and noninfected AEC in monoculture, NKAα1 abundance was rescued to baseline levels at 24h pi, whereas Trail–/– AEC still displayed a strong decrease in NKAα1 protein abundance (Figure 4E). Moreover, transfer of conditioned media from either PBS-treated or PR8-infected AEC to Ifnar–/– AEC did not reduce NKAα1 plasma membrane abundance (Figure 4F), suggesting that in absence of macrophages, NKAα1 surface downregulation was solely dependent on signaling through type I IFN. We next performed coculture infections using AEC and BMM from WT, Ifnar–/–, Trail–/–, or Dr5–/– (TRAIL receptor) mice and assessed NKAα1 plasma membrane abundance after IAV infection. These...
resulted in partial decrease of plasma membrane NKAα1 (Figure 4G).

Addition of recombinant human IFNα and/or recombinant human TRAIL to cultured hAEC reduced their ability for vectorial water transport as measured by changes in FITC-dextran concentrations in the apical versus basal medium of confluent hAEC to a similar extent as IAV infection (Figure 4H), whereas vectorial water transport of hAEC cocultured with primary human AM was significantly increased in the presence of neutralizing antibodies directed against IFNα and TRAIL after IAV infection (Figure 4I). Together, these data indicate that IAV-induced downregulation of NKAα1 and reduced ability for fluid transport depend on a signaling network between AEC and macrophages, involving epithelial type I IFN– and IFN-dependent macrophage TRAIL.

The TRAIL-mediated reduction of NKAα1 levels is induced independently of epithelial cell apoptosis. Previous work on the role of macrophage-released TRAIL during IAV infection demonstrated its contribution to extrinsic AEC apoptosis induction and to loss of barrier function (4, 16). To exclude that the observed effects of TRAIL on NKAα1 expression levels in noninfected AEC were caused by TRAIL-mediated induction of apoptosis, we inhibited activation of the effector caspase-3 by addition of the specific inhibitor Z-DEVD (25, 26) after IAV infection. In vitro and in vivo treatment with Z-DEVD resulted in significantly reduced levels of apoptotic alveolar epithelial cells after IAV infection for 24 hours and 7 days, respectively (Supplemental Figure 4, A and C). Moreover, application of Z-DEVD at d5 and d6 pi in vivo resulted in reduced activation of caspase-3 in AEC at d7 pi (Supplemental Figure 4D). However, NKAα1 plasma membrane expression was not significantly altered after IAV infection by inhibition of caspase-3 activation, both in vitro as well as in vivo (Supplemental Figure 4, B and E).

studies revealed that lack of IFNAR signaling in AEC, together with lack of TRAIL in BMM, fully prevented NKAα1 downregulation in AEC. Partial blockade of the IFN-TRAIL signaling loop by combination of Ifnar−/− AEC with WT BMM (allowing BMM TRAIL action), or of either WT AEC with Tra1−/− BMM or Dr5−/− AEC with WT BMM (allowing AEC type I IFN signaling), correspondingly resulted in reduced activation of caspase-3 in AEC at d7 pi (Supplemental Figure 4D). However, NKAα1 plasma membrane expression was not significantly altered after IAV infection by inhibition of caspase-3 activation, both in vitro as well as in vivo (Supplemental Figure 4, B and E). Accordingly, AFC rates were reduced to similar levels at d7 pi in IAV-infected mice or IAV-infected mice
The IFN/TRAIL-dependent reduction of NKAα1 plasma membrane abundance is mediated via AMPK. Given that Na,K-ATPase endocytosis was reported to be mediated by activation of AMP-activated protein kinase (AMPK) during lung injury–associated treated with Z-DEVD (Supplemental Figure 4F). Together, these data demonstrate that TRAIL-induced reduction of plasma membrane–expressed NKAα1, and its effects on net AFC are mediated independently of the induction of alveolar epithelial apoptosis.

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Figure 4. PR8-induced loss of Na,K-ATPase surface expression is dependent on an IFN-TRAIL signaling loop involving epithelial IFNα and type I IFN–induced macrophage TRAIL. (A and B) IFNα (A) and TRAIL (B) concentrations of AEC mono- and coculture supernatant 24h pi quantified by ELISA. mAEC were monocultured and inoculated in vitro with PBS (ctrl) or PR8 (AEC) or cocultured with BMM without infection (ctrl), with infection of only macrophages (BMM), or with infection of both cell types (AEC/BMM). (C) TRAIL mRNA expression quantified by qPCR in BMM isolated from WT or Ifnar−/− mice noninfected (ctrl) or PR8-infected (IAV) at 16h pi. (D–G) NKAα1 relative MFI on of mAEC treated with 25U/ml mouse rIFNα or/and 100 pg/ml mouse rTRAIL for 16 hours (D); mAEC derived from Ifnar−/− or Trail−/− mice inoculated with PBS (ctrl) or PR8 (IAV) 24h pi (E); mAEC derived from Ifnar−/− mice inoculated for 2 hours with conditioned media from PBS (ctrl) or PR8 (IAV) WT AEC infected for 16 hours (F); and mAEC derived from WT, Ifnar−/−, or Dr5−/− mice cocultured with BMM from WT or Trail−/− mice and infection of none (ctrl) or both cell types (AEC/BMM) in vitro (G). Value of control conditions were normalized to 1. (H and I) Vectorial water transport of hAEC monoculture 6 hours after inoculation with PBS (ctrl), PR8 (IAV), rIFNα (150 pg/ml), and/or rTRAIL (100 pg/ml) or in coculture with human AM without additional treatment (−), in presence of neutralizing antibodies against human IFNα (0.5 μg/ml) and human TRAIL (0.1 μg/ml) (anti-IFNα + anti-TRAIL) or the respective IgG controls (IgG). hAEC maintained barrier integrity in all assay conditions (Supplemental Figure 9). Bar graphs represent means ±SEM of 4–6 independent experiments (A, B, and G–I), 3 independent experiments (C), and 6–8 independent experiments (D). Statistical significance was analyzed by Student’s t test (F) or 1-way ANOVA and post-hoc Tukey (A–E and G–I). *P < 0.05; **P < 0.01; ***P < 0.005.
epithelial dysfunction (27, 28), we investigated if AMPK was involved in regulation of NKA1 abundance in response to IAV-induced paracrine signaling events. AMPK was indeed activated after IAV infection in AEC monolayers and AEC/BMM cocultures (Figure 5, A and B), as demonstrated by phosphorylation of the AMPK downstream substrate acetyl-CoA carboxylase (pACC) (29). Incubation of AEC with conditioned media of IAV-infected AEC and treatment with rTRAIL or rIFNα were sufficient to induce AMPK activation (Figure 5, C and D). Treatment of AEC with the AMPK activator AICA-Riboside (AICAR) (30, 31) led to decreased plasma membrane NKA1 in noninfected AEC and, importantly, chemical inhibition of AMPK by Compound C (32) restored NKA1 levels in infected and in cocultured AEC in presence of infected BMM (Figure 5, E and F). In addition, adenoviral overexpression of a dominant-negative AMPK (DN-AMPK) in human lung epithelial cells (A549, ATCC) rescued NKA1 expression after IAV infection (Supplemental Figure 5A). We found that chemical inhibition of the AMPK upstream calcium/calmodulin-dependent protein kinase β (CaMKKβ) (33) by STO-609 in mAEC or siRNA knockdown of CaMKKβ in A549 resulted in complete inhibition of AMPK activation and restored surface NKA1 abundance in IAV infection (Figure 5, G and H, and Supplemental Figure 5B). In addition, chelation of intracellular calcium needed for CaMKKβ activation by BAPTA-AM increased NKA1 abundance on mAEC after IAV infection (Figure 5G). Inhibition of another possible upstream kinase of AMPK, the TGFβ activated kinase 1 (TAK1), by (5Z)-7-Oxozeanol (34) caused partial inhibition of AMPK activity without affecting NKA1 expression (Figure 5, G and H). Accordingly, inhibition of Na,K-ATPase function by ouabain or IAV infection decreased, whereas inhibition of AMPK or CaMKKβ prevented the reduction in fluid transport after IAV infection (Figure 5I), demonstrating that IFN/TRAIL-induced loss of membrane NKA1 involved a CaMKKβ/AMPK-dependent pathway.

Consistent with the results from our in vitro infection model, inhibition of the intracellular kinase AMPK by intratracheal instillation and overexpression of adenoviral-expressed DN-AMPK in AEC prior to infection in vivo (Supplemental Figure 6A) prevented the IAV-induced reduction in NKA1 plasma membrane abundance and in AFC in vivo at d2 pi (Figure 6, A and B). Additionally, pharmacological inhibition of AMPK signaling in vivo by application of Compound C at d5 pi significantly increased NKA1 plasma membrane abundance and AFC at d7 after IAV infection (Figure 6, C and D). Neither DN-AMPK instillation nor Compound C treatment impacted on viral clearance (Supplemental Figure 6B and data not shown).

IFN and BMM-released TRAIL contribute to reduced NKA1 plasma membrane abundance and impaired AFC in vivo and are amenable to therapeutic targeting after IAV infection. We next investigated whether Na,K-ATPase levels were also dependent on the IFN/TRAIL signaling network in vivo. Interestingly, Ifnar−/− mice (deficient for both IFNα ligation by the alveolar epithelium and for IFNβ-dependent production of TRAIL by macrophages) were protected from IAV-induced reduction of NKA1 plasma membrane expression at d7 pi (Figure 7A). Consistent with our in vitro findings demonstrating that TRAIL released from BMM mediated the reduction of membrane NKA1 abundance during IAV infection, both Trail−/− mice and mice deficient for C-C chemokine receptor type 2 (Ccr2−/−), which are unable to recruit macrophages from the BM to the lung after IAV-infection (35), were protected from NKA1 loss on distal lung epithelial cells in response to IAV infection when compared with WT animals (Figure 7, A and B). Moreover, we demonstrate, by intrapulmonary transfer of BMM flow-sorted (Supplemental Figure 7) from the lungs of WT or Trail−/− mice at d7 after IAV infection into infected Ccr2−/− mice, that rescue of NKA1 expression in Ccr2−/− mice at d7 pi is lost in the presence of WT BMM but maintained after transfer of Trail−/− BMM (Figure 7B). Accordingly, the IFN/TRAIL-mediated loss of surface NKA1 significantly reduced AFC in WT mice after IAV infection but not in Ifnar−/−, Trail−/−, or Ccr2−/− mice (Figure 7C), which corresponded with decreased edema accumulation as quantified by wet-to-dry lung weight ratios in Ifnar−/− and Trail−/− mice (Supplemental Figure 8). Additionally, fluid clearance was completely restored in Ccr2−/− mice that were supplemented with Trail−/− BMM but not in Ccr2−/− mice receiving WT BMM (Figure 7D).

Discussion

Influenza virus pneumonia is characterized by infection of airway and alveolar epithelial cells and can rapidly progress to ARDS with dismal clinical outcome (1, 3). A functional impairment of the alveolar epithelial barrier after IAV infection results in pulmonary edema accumulation and impaired fluid reabsorption, which is closely correlated to mortality in ARDS patients (6, 7). Here, we report a mechanism by which a paracrine communication network between different populations of infected and adjacent cells in the alveolus downregulates alveolar epithelial Na,K-ATPase and thus inhibits AFC in response to IAV infection.

Our data demonstrate that the plasma membrane NKA1 protein abundance is reduced after IAV infection, indicating that the key function of Na,K-ATPase — active sodium transport out of the cell — is impaired (11, 12). Vectorial sodium and fluid clearance is a coordinated process that requires the electrochemical gradient generated by the basolateral Na,K-ATPase, thereby constituting the driving force of edema reabsorption. Sodium and water transport can also be regulated by the apically located ENaC and by chloride channels such as the cystic fibrosis transmembrane receptor (CFTR). In previous work, investigators reported that the levels of ENaC and CFTR were reduced in response to viral infection via a direct interaction with the viral matrix protein (36, 37). In contrast, we found that plasma membrane Na,K-ATPase protein abundance was disproportionately...
The type I IFNs play a complex role in the immune response to IAV infection. They are rapidly produced by AECs after viral infection to activate antiviral transcriptional programs in both epithelial and immune cells in the lung that are important for viral clearance. Furthermore, they may play a role in limiting the severity of the immune response via the production of IL-10 (38–40). Accordingly, IFNs have been suggested as potential therapeutic strategies to promote anti-IAV host defenses, particularly when applied as prophylactic treatment or early after onset of the infection (41, 42). Our data, however, indicate that reduced in neighboring, noninfected epithelial cells, suggesting that NKAα1 reduction is not mediated by direct interaction with a viral component but relies on paracrine signals released from the infected epithelial cells and macrophages. These findings highlight the need to understand influenza A infection in the context of integrated signaling networks involving different cell populations in the lung. Using a coculture system, we demonstrate the importance of epithelial IFNα and IFN-induced macrophage TRAIL, which downregulated the AEC NKAα1 both in the presence and absence of IAV infection.

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Figure 5. IAV-induced loss of NKAα1 surface expression is mediated by activation of AMPK. (A–D) Representative Western blots of n = 3–4 independent experiments of AMPK (62 kDa) and the AMPK substrate pACC (280 kDa) mAEC treated with PBS (ctrl) or PR8 (IAV) (A) for 24 hours. (B) Cocultures of murine AEC and BMM were left uninfected (ctrl); only macrophages were infected (BMM) or both cell types were infected (AEC/BMM). (C) mAEC were treated with conditioned media (CM) of 16 hours PR8-infected (IAV) or PBS-treated (ctrl) mAEC for 2 hours. (D) mAEC treated with the AICAR (1 mM), rTRAIL (100 g/ml), or rIFNα (25 U/ml) for 16 hours. (E–G) Relative MFI of NKAα1 on mAEC inoculated with PBS (ctrl) or PR8 (IAV) for 24 hours without additional treatment or in presence of 1 mM AICAR or 20 μM Compound C (E); mAEC cocultured with BMM without infection (ctrl), with infection of only macrophages (BMM), or with infection of both cell types (AEC/BMM) without additional treatment or in presence of 20 μM Compound C (F); mAEC inoculated with PBS (ctrl) or PR8 (IAV) and incubated for 24 hours without additional treatment or in presence 0.1 μM TAK1 inhibitor (5Z)-7-Oxozeanol or the 0.5 μM CaMKKβ inhibitor STO-609 or 10 μM of the Ca2+ chelator BAPTA-AM (G). (H) Representative Western blot of n = 3–4 independent experiments of AMPK and pACC from 24 hours PR8-infected mAEC without additional treatment or in presence of 20 μM Compound C, 0.1 μM (5Z)-7-Oxozeanol, or the 0.5 μM STO-609 and schematic depiction of the used inhibitors. (I) Vectorial water transport of confluent hAEC culture 6 hours after inoculation with PBS (ctrl) or PR8 (IAV) without additional treatment (–), in the presence of 25 μM Ouabain, 10 μM Compound C, or 0.5 μM STO-609. Bar graphs represent means ±SEM of 4–6 independent experiments. Values of PBS-treated control conditions were normalized to 1. Statistical significance was analyzed by 1-way ANOVA and post-hoc Tukey. *P < 0.05; **P < 0.01; ***P < 0.005.
at later time points, when the infection has spread to the alveolar compartment and progresses to lung injury, IFNs may also have a detrimental effect, evidenced by imbalanced inflammation and loss of crucial AEC functions such as edema clearance.

An important source of IFN and other proinflammatory cytokines in IAV-induced innate immune responses are AM- and CCR2-recruited BMM (43–45). Tissue resident AM play key roles in sensing viral infection and activating the initial innate and later adaptive immune responses to IAV infection (46). In concert with the macrophage populations recruited upon infection, they establish a proinflammatory environment by production and release of mediators such as IFN, leading to enhanced viral clearance and better disease outcomes (43, 47). However, it is well recognized that excessive production of cytokines during IAV infection contributes to lung injury, which impacts severity and outcome (48, 49), as shown for highly pathogenic avian influenza H5N1 or the pandemic 2009 H1N1 virus (50, 51). In particular, resident and recruited macrophages have been attributed as key roles in amplifying lung injury after IAV infection, worsening IAV-induced lung inflammation, lung injury severity, and mortality and hampering resolution of inflammation (4, 16), as inhibition of AEC apoptosis by application of Z-DEVD did not impact on NKAα1 reduction after IAV infection. Activation of the metabolic sensor/regulator AMPK, under normal conditions, upregulates ATP generating and downregulates ATP-consuming mechanisms to promote cellular survival during metabolic stress (34, 55, 56). Of note, Herrero-Martín et al. demonstrated that rTRAIL was able to activate AMPK in immortalized epithelial cells and, thus, promote autophagy, a cytoprotective mechanism rendering cells more resistant to deleterious challenges (57), highlighting the complexity of the IFN/TRAIL network in cellular injury and protection in response to cellular stress.

The Na,K-ATPase has been well documented to be targeted during metabolic stress (34, 55, 56). Of note, Herrero-Martín et al. demonstrated that rTRAIL was able to activate AMPK in immortalized epithelial cells and, thus, promote autophagy, a cytoprotective mechanism rendering cells more resistant to deleterious challenges (57), highlighting the complexity of the IFN/TRAIL network in cellular injury and protection in response to cellular stress.

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The Na,K-ATPase has been well documented to be targeted during metabolic stress (34, 55, 56). Of note, Herrero-Martín et al. demonstrated that rTRAIL was able to activate AMPK in immortalized epithelial cells and, thus, promote autophagy, a cytoprotective mechanism rendering cells more resistant to deleterious challenges (57), highlighting the complexity of the IFN/TRAIL network in cellular injury and protection in response to cellular stress.

Figure 6. IAV-induced loss of NKAα1 surface expression is mediated by AMPK in vivo. (A and B) WT mice were inoculated with PBS (ctrl), Ad-Null, or Ad–DN AMPK for d7 prior to infection with PBS or IAV (Udorn) at d2 pi. (A) Flow cytometric analysis of NKAα1 subunit expression on AEC. (B) In vivo measurements of AFC rates over a time interval of 30 minutes. (C and D) WT mice were inoculated with PBS (ctrl) or PR8 (IAV) and treated with 20 mg/kg Compound C i.p. at d5 pi. Mice were sacrificed at d7 pi. (C) Flow cytometric analysis of NKAα1 subunit expression on AEC. (D) In vivo measurements of AFC rates over a time interval of 30 minutes. Graphs show means ± SEM of n = 4–5 (A), 4–3 (B), 4–9 (C), and 4–6 (D) independent experiments. Data set depicting control conditions in C and D are identical to Figure 1C (AFC) and Figure 3A (NKAα1 expression) and were included for better comparison between experimental conditions. Nonbracketed asterisk indicate statistical significance to control conditions. Statistical significance was analyzed by 1-way ANOVA and post-hoc Tukey. **P < 0.01; ***P < 0.005.
an adverse effect on viral replication (59). Therefore, the resulting changes in intracellular sodium or potassium might interfere with viral signaling, providing an antiviral defense strategy for the infected cell. Alternatively, reduced Na,K-ATPase function might represent activation of a stress response pathway, perhaps secondary to changes in intracellular calcium, that reduces the susceptibility of neighboring epithelial cells to viral infection (59). Consistent with this hypothesis, we found that signaling through IFNAR and DR5 by IFN and TRAIL, respectively, activates AMPK via a pathway that requires CaMKKβ, which is in turn activated by increases in intracellular calcium (58). While the links between calcium signaling and the AMPK during influenza A infection have not been completely elucidated, previous studies have shown that, in hypoxic conditions, an influx of extracellular calcium through stromal interaction molecule 1–activated CRAC channels (57) can lead to CaMKKβ- and AMPK-mediated endocytosis of Na,K-ATPase (27). Wang et al. demonstrated that the calcium-dependent tyrosine kinase Pyk2 interacts with the IFNAR-associated tyrosine kinases Jak1/Tyk2 (60), providing a possible link to CaMKKβ activation.

Our in vitro studies provide complementary lines of evidence demonstrating an additive role for epithelial-produced IFNa- and IFN-dependent macrophage TRAIL in the downregulation of Na,K-ATPase plasma membrane protein abundance during IAV infection. Concordant with our in vitro findings, Ifnar−/− mice deficient in the IFNa/β receptor, and thus compromised in both epithelial IFN crosstalk and IFN-dependent induction of macrophage TRAIL expression, displayed less reduction of Na,K-ATPase and were less impaired in AFC after IAV challenge. Trail−/− mice and mice lacking BMM recruitment from the circulation (Ccr2−/−) were similarly protected from IAV-induced epithelial Na,K-ATPase downregulation. Of course, we cannot fully exclude that additional factors (e.g., infection-induced activation of intracellular pathways) affecting Na,K-ATPase function rather than mere surface expression (e.g., ion or ATP affinity or channel open probability) might have further influence on net AFC in our different in vivo models, as we observe substantially but not fully restored AFC rates ranging between similar levels in all gene-deficient mice applied (Ifnar−/−, Trail−/−, and Ccr2−/−).

Further investigation on the effect of TRAIL released from recruited BMM after infection was performed via pulmonary transfer of WT but not Trail−/− BMM into Ccr2−/− mice after IAV infection, which led to decreased NKAα1 abundance and AFC similar to the decrease observed in IAV-infected WT mice and demonstrated the crucial role for TRAIL-expressing macrophages in the inhibition of AFC in vivo. We acknowledge that, in these studies, we did not address effects on overall survival, nor the consequences of TRAIL-deficiency on virus clearance, but we could importantly show that therapeutic blockade of the IFN/TRAIL network by neutralizing antibodies in vivo proved to increase NKAα1 expression and AFC after IAV infection. Considering that anti-TRAIL treatment was recently demonstrated to elicit profound effects in attenuating IAV-induced pneumococcal superinfection (61), we suggest that therapeutic targeting of the here described signaling axis will not only ameliorate AFC after IAV infection, but will furthermore protect from TRAIL-mediated lung injury and complications thereof. We anticipate that targeting particularly macrophage TRAIL alone or together with epithelial
AMPK downstream signaling might be beneficial with respect to alveolar barrier protection and edema reabsorption, without compromising antiviral host defense.

In conclusion, we describe a paracrine pathway in the IAV-infected lung that links macrophage and epithelial signaling to the downregulation of the Na,K-ATPase in the non-infected fraction of AECs. This decrease is mediated by AEC-released IFNα directly and is strongly amplified through IFNα/β-induced release of TRAIL from recruited macrophages, resulting in inhibition of lung edema clearance (Figure 9). A timely modulation of this pathway might represent a novel strategy to improve fluid reabsorption and, thus, outcomes in IAV-induced lung injury and ARDS.

Methods

Supplemental Methods are available online with this article.

Mice. WT C57BL/6 mice were purchased from Charles River Laboratories. Trail−/− mice (62) were provided by AMGen. U. Kalinke (Paul-Ehrlich Institute, Erlangen, Germany) provided Ifnar−/− mice (63). Dr5−/− mice were a gift from T. Mak (Campbell Family Institute for Breast Cancer Research, University of Toronto, Toronto, Ontario, Canada) (64). Ccr2−/− mice were obtained from W. Kuziel (University of North Carolina Medical School, Chapel Hill, North Carolina, USA; ref. 65). All transgenic lines were backcrossed to the C57BL/6 background. Mice were housed under pathogen-free conditions.

Cell isolation and culture. Primary mAEC or hAEC were isolated as previously described (16, 66), seeded on 4 μm–pore size transwells (Corning Inc.), and cultured for 3 and 5 days, respectively, prior to experiments. AEC suspensions with a purity >90% were used for further experiments. Murine resident AMs were isolated by bronchoalveolar lavage as described (16) and cultured for 2 hours before infection. Human resident AMs were gained from bronchoalveolar lavage fluid (BALF) of patients who did not show any accumulation of pappenheim-stained cytopsins. BMM were isolated from femurs and tibias and cultured in the presence of mouse recombinant granulocyte/macrophage-CSF (GM-CSF) at 25 ng/ml for 10 days.

Infection protocols. Cells were infected with PR8 at a multiplicity of infection (MOI) of 0.1 as previously described (16). For coculture experiments, bottom-seeded AM or BMM were combined with AEC seeded on transwells after infection of only macrophages or of both cell types with PR8. Of note, an abortive replication cycle occurred in AM and BMM, as described in Högnér et al. (16), and ensured that in the AM/BMM infection–only conditions, no virus was transferred to the epithelial cell layer excluding AEC infection. For in vitro experiments, recombinant murine TRAIL (R&D Systems, 100 pg/ml), recombinant human GM-CSF (R&D Systems, 25 ng/ml), recombinant mouse IFNα and -β (PBL Interferon Source, 25 U/ml; R&D Systems, 10–100 ng/ml), recombinant human IFNα (R&D Systems, 150 pg/ml), recombinant human TRAIL (R&D Systems, 100 pg/ml), anti–human IFNα neutralizing antibody (R&D Systems, 0.5 μg/ml), and anti–human TRAIL neutralizing antibody (R&D Systems, 0.1 μg/ml), and the following chemical inhibitors were used: AICAR, AMPK inhibitor Compound C, (5Z)-7-Oxozeanol (Curvularia sp), STO-609 (all Calbiochem) and BAPTA-AM (Invitrogen). Conditioned media of 16-h-infected mAEC cultured in 6-well plates was transferred to uninfected mAEC.
cultured in 12-well plates and incubated for 2 hours. For in vivo infection, mice were intratracheally inoculated with 500 plaque-forming units (pfu) PR8 or 10⁷ pfu A/Udorn/307/1972 (H3N2, also known as Udorn) diluted in 70-μl sterile PBS. Endotracheal delivery of adenovirus was performed at 1 × 10⁹ pfu in 50% surfactant vehicle as described previously (24). Adenoviruses carrying either no additional construct (Ad-Null) — or expressing DN-AMPK, Ad-NKAβ1, or Ad-NKAβ1 — were cloned at ViraQuest Inc. Of note, NKAβ1 — which was shown to be a crucial mediator for correct heterodimerization of the Na,K-ATPase (22, 67) — was coexpressed to ensure presence of sufficient levels of NKAβ1 to support functional Na,K-ATPase formation.

Of note, transduction efficiency was >90% in AEC, as verified by FACS for mCherry in EpCAM⁺ cells (not shown). Neutralizing antibodies against IFNa (10,000 IU/mouse, rabbit polyclonal Ab to mouse IFNa, PBL Interferon Source) were administered intratracheally at d3 pi, and neutralizing antibodies against TRAIL (150 μg/mouse, LEAF anti-mouse CD253, BioLegend) were given i.p. at d3 and d5 pi as established previously (16). Infected mice were monitored 1–3 times per day.

**In vivo measurement of AFC.** The rate of fluid removal from the alveolar airspace was measured by assessment of changes in Evans blue–tagged albumin in an iso-osmolar alveolar instillate over 30 minutes as previously described (24). In short, mice were anesthetized, placed on a heating pad, and maintained in supine position. Body temperature was monitored using an anal probe and maintained at 37°C with heating pad and lamp. After reaching stable anesthesia, mice were cannulated with a shortened 18-gauge i.v. catheter, paralyzed with 2.0 mg/kg pancuronium bromide applied i.p., and ventilated via catheter connected to a mouse respirator (Harvard Apparatus) using a tidal volume of 10 ml/kg (equivalent of 0.2 ml) at a frequency of 160 breaths per minute and 100% of oxygen, as described by Hardiman et al. (68). Iso-osmolar NaCl solution (324 mOsm; 300 μl) containing 5% Evans blue–tagged (0.15 mg/ml) bovine serum albumin were installed via the catheter over a time period of 30 seconds, followed by 0.1 ml of air; then, mice were ventilated over 30 minutes. Evans blue concentrations, directly correlating with the amount of BSA in the solution, were analyzed from the initial instillate and the reaspirated fluid using a microplate reader (Bio-Rad, 620-nm filter) and AFC was calculated as follows: AFC = 1 – (C₀/Cₐ) – [1 – (C/Cₐ)], where C₀ is the protein concentration before instillation and Cₐ is the protein concentration of the sample reaspirated after 30 minutes of ventilation.

**Ex vivo measurement of vectorial water transport.** Primary human AEC were seeded in 0.4 μm–pore size transwell cell culture dishes (Corning Inc.) and cultured until achieving electrochemical resistances of ≥800 Ω/cm² as measured by Millicell-ERS2 device (EMD Millipore). Cells were infected with PR8 at MOI 0.1 or PBS treated for 1 hour at 37°C and then supplied with 3 mg/ml 70 kDa FITC-dextran–labeled (Sigma-Aldrich) cell culture media including selected inhibitors at indicated concentrations. After 6 hours of incubation at 37°C, apical and basal media were analyzed for FITC-dextran concentration (FLₓ 800, Bio-Tek). Cells were analyzed microscopically for bound FITC-dextran without apparent differences in the treatment groups. Vectorial water transport was calculated by changes in FITC-dextran concentration between apical (Cₐ) and basal (Cₐ) media in comparison with starting conditions (C₀): [1 – (Cₐ/C₀)] – [1 – (C/Cₐ)].

**Histology.** Lungs were clipped at the trachea, perfused with 4% paraformaldehyde (PFA), removed, and fixed for 24 hours in 4% PFA. Lungs were embedded in Paraffin (Leica ASP200S), cut into 5-μm thick sections, and either stained with NKAβ1 (clone 40H9) and phospho-ACC (clone S79, both Cell Signaling Technology), β-actin (clone Poly6221, BioLegend), GAPDH (clone 14C10, Cell Signaling Technology), secondary mouse or rabbit HRP (clones 7076 and 7074, Cell Signaling Technology). Bands were detected...
with the MicroChemi system (DNR Bio-Imaging Systems Ltd.) and quantified using ImageJ software.

Flow cytometry and cell sorting. Multicolor flow cytometry and cell sorting were performed with an LSR Fortessa and BD FACSaria III cell sorter using DIVA software (BD Biosciences) as previously described (16, 70). Cells (1 × 10^6) from AEC cell cultures or derived from lavaged, perfused, and homogenized murine lungs were freshly stained with fluorochrome-labeled antibodies for 15 minutes at 4°C in BD FACs buffer (BD Biosciences). When nonlabeled primary monoclonal antibody was used, a fluorescent-labeled secondary antibody was incubated for 15 minutes at 4°C in FACs buffer. Cells were routinely stained with 7-AAD (Invitrogen) for dead cell exclusion, and were also stained with antibodies to detect CD326 (clone G8.8, BioLegend), CD74 (clone ln-1, BD Biosciences), Podoplanin (clone 8.1., BioLegend), polyclonal IAV (clone ab20841, Abcam), NKAa1, CD45 (clone 30-F11, BioLegend), GR-1 (clone RB6-8C5, BioLegend), Ly6G (clone IA8, BioLegend), SiglecF (clone ES0-2440, BD Biosciences), CD11c (clone N418, BioLegend), CD11b (clone M1/70, BioLegend), secondary goat APC (clone A21447, Invitrogen), and secondary rat PE (clone A10545, Invitrogen). Corresponding isotype antibodies were used as negative controls. For annexin V staining, cells were resuspended in annexin V staining buffer (10 mM HEPES, 140 mM NaCl, and 2.5 mM CaCl$_2$) and incubated with annexin V Alexa Fluor 647 (1:100, Invitrogen) for 15 minutes at 4°C. Data are presented as median fluorescent intensity (MFI) and were normalized to control groups set to 1 for in vitro experiments. NKAa1+ AEC from in vivo–infected mice are given as a percentage of the epithelial (EpCAM+) cell population.

ELISA. Cell culture supernatants were analyzed using commercially available ELISA kits for mouse TRAIL (R&D Systems, detection limit 1.8 pg/ml) and mouse IFN-β (PBL Interferon Source, detection limit 12.5 pg/ml), according to the manufacturer’s instructions.

Quantitative PCR. RNA was isolated using RNeasy Kit (QIAGEN) and cDNA synthesized as described previously (66). Quantitative PCR (qPCR) was performed with SYBR green I (Invitrogen) in the AB StepOnePlus Detection System (Applied Bioscience). β-Actin expression served as normalization control. Data are presented as ddCt (dCt_target - dCt_germ) or fold-change (2^(-ΔΔCt)). The following primers were used: β-actin (forward primer, 5′-ACCCTAAGGCAACGTGA-3′; reverse primer, 5′-CAGAGGCTACAGGGACAGCA-3′); TRAIL (forward primer, 5′-GAAGACCTCAGAAATTGGC-3′; reverse primer, 5′-GACCCGCTCCTCCTGTTA-3′).

Pulmonary transfer of BMM. BMM of WT or Tail+/- mice recruited to the alveolar compartment after IAV challenge with 500 pfu PR8 were obtained by bronchoalveolar lavage on d7 pi after challenge with 500 pfu PR8. BMM were identified by the signature CD45+GR-1-Ly6G CD11b+CD11c+SiglecF+ and flow sorted (Supplemental Figure 5). The purity of sorted BMM (≥95%) was ensured by flow cytometric analysis, as well as by pappenheim stained cyto-  

Statistics. All data are given as mean ± SEM. Statistical significance was analyzed by unpaired 2-tailed Student’s t test or by 1-way ANOVA and post-hoc Tukey (GraphPad Prism 5). A P value less than 0.05 was considered significant. *P < 0.05; **P < 0.01; ***P < 0.005.

Study approval. Animal experiments were approved by the regional authorities of the State of Hesse (Regierungspräsidium Giessen) and by the Institutional Animal Care and Use Committee at Northwestern University. Human lung tissue was obtained from patients who underwent lobectomy after informed written consent. Use of human lung tissue and BLM samples was approved by the University of Giessen Ethics Committee.

Author contributions CP, CS, IV, TW, SP, REM, JL, WS, and SH were involved in study design and concept. Data were acquired by CP, BS, LMN, EL, JB, LF, and SG. GMM and GRSB were involved in the AFC measurements. Data analysis, interpretation, and statistics was conducted by CP, EL, KM, JL, JIS, IV, and SH. CP, CS, JL, JIS, and SH drafted the manuscript.

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Address correspondence to: Susanne Herold, Universities Giessen & Marburg Lung Center, Department of Internal Medicine II, Klinikstr. 33, D-35392 Giessen, Germany. Phone: 49.641.985.42552; E-mail: Susanne.Herold@innere.med.uni-giessen.de.

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